CARE TRANSITIONS NOTEBOOK

Caring for Someone with Memory Loss or Alzheimer’s After a Hospitalization

Permission granted by Alzheimer’s Greater Los Angeles. Adapted from Care Transitions Notebook: Caring for Someone with Memory Loss or Alzheimer’s After a Hospitalization; developed by Alzheimer’s Greater Los Angeles

© 2016 Alzheimer’s Greater Los Angeles
# Table of Contents

Commonly Used Words........................................................................................................1
Stages of Alzheimer’s Disease ..............................................................................................2
Self-Care .............................................................................................................................3
Hospital Discharge Plan....................................................................................................4
Understanding “Baseline”.................................................................................................5
Knowing When to Call the Doctor ..................................................................................6
Medications......................................................................................................................8
Warning Signs of Pain.......................................................................................................9
Keeping the Home Safe ...................................................................................................10
Challenging Behaviors.....................................................................................................11
IDEA!..................................................................................................................................12
Common Challenging Behaviors and Possible Adaptations ............................................14
Being a Healthier Caregiver ............................................................................................21
Personal Goal Checklist..................................................................................................22
North Carolina Resources ...............................................................................................23
Purpose of Care Transitions Notebook

Even a brief stay in the hospital can be very stressful for a person who has Alzheimer's disease or other type of dementia. In addition to being ill, being in an unfamiliar place and dealing with strangers can result in changes in memory, thinking, and behavior. It may take the person several weeks after going home to return to their usual way of thinking and behaving. This notebook was developed to help caregivers understand and plan for care needs of patients with Alzheimer’s disease who have been in the hospital.

Commonly Used Words

Throughout this notebook, you will see some commonly used words. For purposes of this project, they are explained below.

**Care transitions:** the 30 days after a person leaves the hospital and is back at home. During this time, it is important that the person’s needs are coordinated so his/her health and social needs are met.

**Dementia:** a general word that means a person has problems remembering, and problems with other thinking skills that are bad enough to get in the way of day-to-day living. For example, it may be hard for the person to make decisions or pay attention. He/she may have a hard time finding the right words when talking. Dementia is not a specific disease and has many causes. Throughout this Care Transitions Notebook, we talk about “Alzheimer’s disease,” but other diseases can cause problems with memory and thinking, too.

**Alzheimer’s disease:** a disease that destroys memory and functioning (like being able to take care of yourself). Alzheimer’s disease happens slowly and gets worse over time. Alzheimer’s disease is one kind of dementia.

**Delirium:** a condition caused by a medical problem that gets worse; can look like serious confusion and can develop over hours or days; needs to be treated by a doctor.
Stages of Alzheimer’s Disease

Not everyone will have the same signs of Alzheimer’s disease at the same time. Unfortunately, Alzheimer’s does get worse over time. It is important to understand what to expect in the different stages of the disease. The more you know, the better you can prepare.

We usually talk about Alzheimer’s disease having three main stages:

- early stage
- middle stage
- late stage

This Care Transitions Notebook focuses on people in the middle stage of the disease.

<table>
<thead>
<tr>
<th>STAGES OF ALZHEIMER’S DISEASE</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>EARLY STAGE</strong></td>
</tr>
<tr>
<td>Families, friends, and co-workers may notice problems with memory and concentration</td>
</tr>
<tr>
<td>Trouble finding the “right” word and/or remembering names</td>
</tr>
<tr>
<td>Misplacing things</td>
</tr>
<tr>
<td>Trouble organizing and planning</td>
</tr>
<tr>
<td>Person often able to make decisions and plan ahead</td>
</tr>
<tr>
<td>Caregiver is more involved</td>
</tr>
</tbody>
</table>

*In the early stages of the disease, it is important that you include the person with Alzheimer’s disease in decision-making and care planning.*
How Well Can a Person with Alzheimer’s Disease Take Care of Himself/Herself?

In the beginning of the disease, the person with Alzheimer’s can make decisions and take care of himself/herself, but this will change. Eventually, everyone with Alzheimer’s disease will need a person to help them with day-to-day care. We call that person a “caregiver.”

In the middle stage of Alzheimer’s, a caregiver may need to help the person with:

- following hospital discharge instructions
- making sure the person with Alzheimer’s is not left home alone
- taking correct medicines
- bathing, dressing, eating, toileting, and other activities
- making doctor’s appointments and following the doctor’s instructions
Using the Hospital Discharge Plan to Understand Care

Look at the hospital discharge plan and write down the main things you, as a caregiver, need to do to help the person with Alzheimer’s. Remember that someone who has middle stage Alzheimer’s will need help with medications and overall care.

1) 

2) 

3) 

*If you did not get a hospital discharge plan or you lost it, call your doctor to ask for instructions.
Understanding “Baseline”

Baseline is a word that doctors use to describe how a person usually thinks and acts. When someone has Alzheimer’s, his/her thinking and behavior can change, but it is usually a slow change.

What is “USUAL?”

Everyone is different, but you know what the person you are caring for is usually like. When someone has Alzheimer’s, they have some days that are better than others, but overall, they act and think a certain way.

"My grandpa was usually a happy person. He would sing and dance. He never yelled or got angry. When all of a sudden he started yelling at people and saying mean things, I was concerned. It just wasn’t like him. I called the doctor immediately and it turned out that my grandpa had an infection. Once the infection was treated, my grandpa went back to being his usual nice and happy self."

- Anonymous caregiver
Knowing When to Call the Doctor: Warning Signs of Health Issues & Common Causes of Delirium

- People with Alzheimer’s disease may not be able to tell you with words or full sentences that something is wrong. They may not be able to say they are not feeling well.

- You should call the doctor if you see that the person has had a sudden and unusual change in the way he/she is acting or thinking, or has a medical condition that is getting worse. These changes can happen over a period of several days.

- Be prepared to tell the doctor what the person is usually like (before he/she had the sudden and unusual change).

Write here what the person is usually like before the sudden and unusual change:

________________________________________________________________________
________________________________________________________________________
________________________________________________________________________
________________________________________________________________________

- If you think there is a medical emergency, call 911.

Understanding delirium

People with Alzheimer's are at risk for developing delirium.

Delirium is usually caused by an illness or reaction to medication.

Delirium looks like severe confusion and can develop over hours or days. The person may seem “out of it” and then become alert again. The person may also have changes in sleeping patterns, with vivid/intense dreams.

It is important to call the doctor if you see any of these changes.
What are examples of sudden and unusual behaviors that you should look for?

Here are some examples of things that can happen **SUDDENLY:**

- major change in memory or mood
- increase in confusion
- not knowing where he or she is, or what time it is
- cannot pay attention
- angry, hitting, and yelling (becoming aggressive or violent)
- going to the bathroom in his/her pants or wetting the bed
- fever
- seizure
- hallucinations (seeing, hearing, smelling, tasting or feeling something that isn't there)

If you’ve seen a sudden and unusual change that concerns you, write it down and contact the doctor:

---

Write down the name and phone number of the doctor:
Taking Medications

People with Alzheimer’s disease will eventually need help taking their medications. Taking too much of a medication, taking too little, or not following the directions can be dangerous.

You cannot rely on the person with Alzheimer’s disease to take his/her medications; you will need to make sure the medications are taken correctly.

Also, make sure that medications are locked up so they are out of reach.

Why is it important that you assist with medications?

You will help make sure that:

- the right medications are taken
- at the right time
- and the right amount

* When you see the doctor, take all of the medications, vitamins, supplements, and herbs with you.
Warning Signs of Pain

Communication can become challenging when a person has Alzheimer’s disease. The person may not be able to tell you that he or she is in pain. If you think that the person is in pain, call the doctor.

Remember that people with Alzheimer’s disease feel pain just like people who do not have Alzheimer’s. Pain is usually something that can be treated.

What are the possible signs of pain?

- **Physical signs**
  - bruises
  - swelling
  - fever
  - throwing up
  - dry/pale gums
  - sores on the body and in the mouth
  - pale/light skin tone
  - flushed/red skin tone

- **Nonverbal signs**
  - gestures/movements, like holding a part of the body
  - spoken sounds like groans or grunts
  - facial expressions like wincing or grimacing

- **Changes in behavior**
  - increased anxiety
  - increased agitation
  - shouting
  - new sleeping problems

Source: Alzheimer’s Association [http://www.alz.org/care/alzheimers-late-end-stage-caregiving.asp #pain#ixzz352NT7Fx1](http://www.alz.org/care/alzheimers-late-end-stage-caregiving.asp #pain#ixzz352NT7Fx1)
Keeping the Home Safe

Caregivers need to make sure that the home is safe for the person they are caring for. A person with Alzheimer’s disease may have trouble knowing what is dangerous. By helping the person feel more relaxed and less confused at home, you can help stop accidents.

What are things that can help with safety?

**Never leave a person with Alzheimer’s home alone**

- If you need to leave the house, either take the person with you or find someone to stay with him/her while you are gone

**Reduce the risk for falls**

- Keep rooms neat
- Remove small rugs and mats or anything that might slide on the floor
- Keep things off of the floor…cords, books, toys, bags, boxes, etc.
- Use tables and chairs that are stable enough to lean on
- Use a night light at night so the person you are caring for can see where he/she is going
Challenging Behaviors

When a person has Alzheimer’s disease, he/she may have some challenging behaviors. These behaviors may be challenging for the person with Alzheimer’s and/or may be challenging for you.

Some examples of challenging behaviors are:

- getting angry and fighting
- wandering or getting lost
- hallucinations (seeing, hearing, smelling, tasting, or feeling something that isn’t really there)
- paranoia (not trusting other people)

These behaviors tell us that the person with Alzheimer’s needs something, or that something is wrong. These behaviors are not done on purpose; they are part of the disease.

If these behaviors suddenly become worse or are unusual, call the doctor.

“My father would sometimes get really agitated and say things that concerned us. He would think that the house was being bombed when he heard airplanes or helicopters. He was probably thinking back to the time when he was a fighter pilot. We had to find creative ways to comfort him and reassure him that he was safe. Staying calm and speaking gently helped.”

- Daughter
IDEA!

IDEA! is a simple three-step strategy to help you figure out why a challenging behavior is happening and how to deal with it.

**Identify the problem/challenging behavior**

- What is the behavior that is challenging for you to deal with?
  - Be specific. Can you see it?

**Educate Yourself**

**Understand the cause of the behavior**

- **Health:** Is the person taking a new medication, getting sick, or in pain?
- **Environment:** Is it too noisy? Is it too hot? Is the place unfamiliar?
- **Task:** Is the task too hard? Are there too many steps? Is it something new?
- **Communication:** Is it hard for the person to speak or understand?

**Understand the meaning of the behavior to the person**

- Does the person feel like he/she is being treated like a child?
- Are there things that remind the person of something unhappy?
- Does the person feel a sense of insecurity, discomfort, or boredom?

**Adapt**

**Try different things. Pay attention to the person’s feelings. Practice being calm, gentle, and reassuring.**

- **Distract or redirect by**
  - Offering the person something he/she likes to eat
  - Watching a TV show or listening to music
  - Asking the person for his/her help with a simple activity
  - Leading the person to a different room

- **Address the cause or triggers of the behavior**
  - Keep tasks and activities simple
  - Keep the home as quiet and calm as possible
  - Speak slowly and gently/try not to say too much at one time
  - Don’t argue/try to comfort the person
  - Find meaningful, simple activities so the person isn’t bored
Try Using *IDEA*!

Write down a behavior that has been challenging for you to deal with. Use *IDEA* to break it down and figure out some possible solutions.

**IDentify the problem**
The challenging behavior is ________________________________

**Educate Yourself**

**Understand the cause of the behavior**
When do you see this behavior happening? ________________________________

What are the things that seem to trigger the behavior? ________________________________

**Understand the meaning of the behavior**
What might this behavior tell you about how the person is feeling?

**Adapt**
What can you try doing differently? ________________________________

—

“*My mother would scream every time we tried to bathe her. When I put myself in her shoes, I realized that it was cold in the bathroom and she was uncomfortable getting undressed. As soon as I made the temperature in the bathroom warmer and gave her extra towels to cover up, she stopped yelling. It was important to understand what was *causing* the yelling so I could make some changes to the environment.*”

- Daughter and caregiver
## Common Challenging Behaviors and Possible Adaptations (Solutions)

<table>
<thead>
<tr>
<th>Behavior</th>
<th>Possible Adaptations (Solutions)</th>
</tr>
</thead>
</table>
| **Agitation/Combativeness** | - Respond in a calm way; use a gentle voice  
- Offer encouragement  
- Use short, simple sentences  
- Make tasks more simple by breaking things down step-by-step  
- Give the person enough time to respond  
- Approach the person slowly and from the front  
- Avoid fighting with the person or correcting them  
- Distract the person with another enjoyable activity  
- Go for a walk  
- Find a quiet place to sit and relax  
- Find a comforting object (like a stuffed animal)  
- Don’t expect the person to do more than he/she can do  
- Keep the home calm, quiet, and clutter free |
<table>
<thead>
<tr>
<th>Behavior</th>
<th>Possible Adaptations (Solutions)</th>
</tr>
</thead>
</table>
| **Dressing** | - Give the person extra time  
- Don’t act like you are in a hurry  
- Limit the person’s choices to two outfits; let him/her decide which outfit to wear  
- Lay the person’s clothes out in the order needed to put them on  
- Talk the person through getting dressed using short, simple, one-step instructions  
- If the person loves a certain outfit and refuses to wear anything else, buy several outfits that look the same  
- Use pants with elastic waistbands and pullover tops to make getting dressed easier |
| **Eating** | - Give the person lots of time to eat; do not rush  
- Serve meals at the same time every day  
- Serve foods with different colors and textures  
- Use plain-colored dishes (without patterns or dark colors) so the person can see the food on the plate  
- Use a shallow bowl with a lip on it if the person keeps pushing food off the plate  
- Put only the needed utensils next to the plate  
- Try offering one food at a time  
- Help the person with eating if he/she is having a hard time, but let the person do as much as possible  
- Try finger foods |
<table>
<thead>
<tr>
<th>Behavior</th>
<th>Possible Adaptations (Solutions)</th>
</tr>
</thead>
</table>
| **Hallucinations**  
(seeing, hearing, smelling, tasting, or feeling something that isn’t really there) | - Don’t argue with the person that something is not real (because it is real to the person)  
- Comfort the person; try saying, “I am here and I will take care of you”  
- Use a calm and gentle tone of voice  
- Try to change locations if something nearby is triggering the hallucinations  
- Find a relaxing and enjoyable activity  
- Go for a walk  
- Cover mirrors and windows if the person doesn’t know who is in the mirror or window  
- Turn off the TV if it is confusing  
- Turn on lights; use a nightlight  
- If hallucinations start suddenly, call the doctor |
| **Pacing**       
(moving back and forth) | - Walk with the person  
- Hold the person’s hand  
- Tell the person he/she is safe and loved  
- Offer the person a snack or a comforting object to hold in his/her hand as he/she walks  
- Keep walkways clear so the person doesn’t fall  
- Try to distract. For example, ask the person to look at a magazine with you or to help you with a puzzle |
<table>
<thead>
<tr>
<th>Behavior</th>
<th>Possible Adaptations (Solutions)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Paranoia</strong></td>
<td>- Don’t be offended if the person accuses you of something (like stealing)</td>
</tr>
<tr>
<td>(not trusting other people)</td>
<td>- Don’t argue or try to convince the person</td>
</tr>
<tr>
<td></td>
<td>- Give simple answers</td>
</tr>
<tr>
<td></td>
<td>- Distract with an enjoyable activity</td>
</tr>
<tr>
<td></td>
<td>- If the person is often looking for a specific item, have extras available (for example, if the person is always looking for his/her wallet, buy two)</td>
</tr>
<tr>
<td><strong>Repetition</strong></td>
<td>- Look for the reason behind the repeating</td>
</tr>
<tr>
<td>(saying or doing the same thing over and over)</td>
<td>- Focus on the emotion, not the behavior (think about how the person is feeling)</td>
</tr>
<tr>
<td></td>
<td>- Turn the action or behavior into an activity (for example, if the person is rubbing his/her hand across the table, provide a cloth and ask for help with cleaning)</td>
</tr>
<tr>
<td></td>
<td>- Stay calm and be patient</td>
</tr>
<tr>
<td></td>
<td>- Use a gentle and calm voice</td>
</tr>
<tr>
<td></td>
<td>- Don’t argue or try using logic</td>
</tr>
<tr>
<td></td>
<td>- Provide the person with the answer he/she is looking for</td>
</tr>
<tr>
<td></td>
<td>- Distract with an enjoyable activity</td>
</tr>
<tr>
<td></td>
<td>- Accept the behavior and try to work with it (as long as it is not dangerous)</td>
</tr>
<tr>
<td>Behavior</td>
<td>Possible Adaptations (Solutions)</td>
</tr>
<tr>
<td>----------</td>
<td>--------------------------------</td>
</tr>
<tr>
<td><strong>Toileting</strong></td>
<td></td>
</tr>
</tbody>
</table>
- Mark the bathroom clearly with a sign that says “bathroom” and with a picture of a toilet  
- Watch for signs like fidgeting with clothing, pacing, or unusual sounds or faces  
- Walk with the person to the bathroom every 2 to 3 hours and do not wait for the person to ask; don’t make a big deal out of it; say in a happy, calm, and gentle voice, “It’s time for us to go to the bathroom”  
- Carry extra toileting supplies with you when you are away from home  
- Leave on a nightlight in the bathroom  
- Keep the person’s dress as simple as possible (choose easy-to-remove and easy-to-clean styles such as sweat pants with elastic waistbands) |
<table>
<thead>
<tr>
<th>Behavior</th>
<th>Possible Adaptations (Solutions)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sundowning</td>
<td>▪ Turn on more lights in the afternoon and evening</td>
</tr>
<tr>
<td>(difficult behaviors in</td>
<td>▪ Make afternoon and evening hours less busy (schedule appointments, trips, and activities</td>
</tr>
<tr>
<td>the late afternoon and</td>
<td>earlier in the day)</td>
</tr>
<tr>
<td>evening)</td>
<td>▪ Help the person use up extra energy with exercise</td>
</tr>
<tr>
<td></td>
<td>▪ Reduce foods and drinks with caffeine</td>
</tr>
<tr>
<td></td>
<td>▪ Give the person an early dinner or late afternoon snack</td>
</tr>
<tr>
<td></td>
<td>▪ Try to avoid or limit naps during the day</td>
</tr>
<tr>
<td></td>
<td>▪ Lower the noise level</td>
</tr>
<tr>
<td></td>
<td>▪ Close the blinds or curtains</td>
</tr>
<tr>
<td></td>
<td>▪ Tell the person where he/she is and that he/she is safe</td>
</tr>
<tr>
<td></td>
<td>▪ Tell the person you are not leaving</td>
</tr>
<tr>
<td></td>
<td>▪ Use a happy, calm, and gentle voice</td>
</tr>
<tr>
<td>Behavior</td>
<td>Possible Adaptations (Solutions)</td>
</tr>
<tr>
<td>--------------------------</td>
<td>------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Wandering/ getting lost</td>
<td>- Register person for Medic Alert®+Alzheimer’s Safe Return®</td>
</tr>
<tr>
<td></td>
<td>- Make sure that the person wears the Medic Alert® ID bracelet</td>
</tr>
<tr>
<td></td>
<td>- Keep a recent photograph of the person to help police if he/she should become lost</td>
</tr>
<tr>
<td></td>
<td>- Keep all doors locked</td>
</tr>
<tr>
<td></td>
<td>- Place safety latches up high and down low on doors</td>
</tr>
<tr>
<td></td>
<td>- Place cloth of the same color of the door over door knobs, or paint the doors and doorknobs</td>
</tr>
<tr>
<td></td>
<td>the same color as the walls</td>
</tr>
<tr>
<td></td>
<td>- Make sure the person gets enough exercise and sleep; staying active may help</td>
</tr>
<tr>
<td></td>
<td>- Let the person do chores, such as folding clothes or helping with dinner</td>
</tr>
</tbody>
</table>
Being a Healthier Caregiver

The only way you will be able to continue helping someone with Alzheimer’s is to make sure that you are also taking care of yourself. Think about your physical, mental, and emotional health.

Here are some tips on how to be a healthier caregiver:

- Find time for yourself and do the things you like to do
- Get enough rest, eat right, exercise, and visit your doctor
- Manage your stress level
- Have a backup plan in case something unexpected happens to you
- Plan each day, but remember that you will need to be flexible too
- Be realistic
- Pat yourself on the back for the good job you are doing
- Become an educated caregiver; know what resources are available, get help, and find support
- Reach out for help and support - talk to others about how you feel - join a support group

10 signs of caregiver stress

If you are experiencing the following signs, contact your doctor:

- Denial
- Anger
- Removing yourself from friends/family and activities
- Anxiety/nervousness
- Depression
- Exhaustion (very tired)
- Not able to sleep
- Irritability
- Cannot concentrate
- Problems with physical health

Source: Alzheimer’s Association. www.alz.org
Personal Goal Checklist

You may not be able to work on personal goals right away, but try to pick one goal to start thinking about. Remember that taking care of yourself will also help the person you are caring for.

<table>
<thead>
<tr>
<th>Personal Goals for Caregiver</th>
</tr>
</thead>
<tbody>
<tr>
<td>☐ Who can relieve me for an hour or two so I can take a break?</td>
</tr>
<tr>
<td>☐ Who can spend the night with the person I am caring for so I can get a good night’s rest?</td>
</tr>
<tr>
<td>☐ Who can I call if I am feeling overwhelmed?</td>
</tr>
</tbody>
</table>

Write down what you need to do to start working on this personal goal:

______________________________________________________________

______________________________________________________________

______________________________________________________________

______________________________________________________________

______________________________________________________________
North Carolina Resources for Alzheimer’s Disease and Related Dementias

Alzheimer’s Association National Office provides a Helpline and website for information about Alzheimer’s Disease. The Helpline (1-800-272-3900) is available 24 hours per day, 7 days per week, and provides translation services in more than 200 languages. There are two chapters of the Alzheimer’s Association in North Carolina: Alzheimer’s Association-Eastern North Carolina and Alzheimer’s Association-Western North Carolina. For information about resources and support groups in your local area, go to http://www.alz.org/northcarolina.

MediAlert® + Alzheimer's Association Safe Return® is a nationwide emergency response service for individuals with Alzheimer's or a related dementia who wander or have a medical emergency. Assistance is available 24 hours a day, 365 days a year. The program provides an ID bracelet or pendant to be worn by the individual with dementia. There are one-time enrollment and annual renewal fees. To sign up for this service, call 1-888-572-8566. Go to http://www.alz.org/care/dementia-medic-alert-safe-return.asp#ixzz4h9reByjZ for additional information.

Alzheimer’s North Carolina offers information and referral and educational programs and services, including support groups and a toll-free Helpline, for individuals with Alzheimer’s disease and related disorders and their caregivers. Alzheimer’s NC is not affiliated with the national Alzheimer’s Association. For additional information, or to locate a support group in your area, go to http://www.alznc.org. The toll-free number is 1-800-228-8738.

Area Agencies on Aging (AAAs) provide information on a range of available services and assistance for older adults, adults with disabilities, and their caregivers, including:
- information on available services in your area (including meals, senior centers, adult day care, etc.);
- assistance in gaining access to services;
- individual counseling, support groups and caregiver training;
- respite care; and
- supplemental services of various kinds
Go to https://www.ncdhhs.gov/assistance/adult-services/area-agencies-on-aging to contact your region’s AAA.

BenefitsCheckUp® is a free, comprehensive online tool that connects older adults with benefits they may qualify for. Go to https://www.benefitscheckup.org/#/.

Duke Family Support Program offers support to families caring for someone with Alzheimer’s or other memory disorders. The program offers these services to all NC residents at no charge:
- tips on caring for people with memory disorders;
- an information packet on Alzheimer’s and related diseases;
- phone or email help with care decisions, evaluating assisted living or nursing facility care, and coping strategies;
- help selecting support groups, education programs, websites or books; and
- The Caregiver newsletter (twice yearly) and Triangle Area E-News (monthly).
Call 1-800-646-2028 or go to http://www.dukefamilysupport.org.

Medicare.gov website ([https://www.medicare.gov/](https://www.medicare.gov/)) provides comparison information for hospitals, skilled nursing facilities, home health agencies and other providers.

**Memory Cafes** are welcoming places for individuals with Alzheimer’s or any type of dementia or other brain disorder and their caregivers. They are safe and comfortable spaces where caregivers and their loved ones can socialize, listen to music, play games and enjoy other activities together. For a list of Memory Café locations in North Carolina, go to: [http://www.memorycafedirectory.com/memory-cafes-in-north-carolina/](http://www.memorycafedirectory.com/memory-cafes-in-north-carolina/)

**North Carolina Seniors’ Health Insurance Information Program (NC SHIIP)** offers a toll-free line (1-855-408-1212) and local counseling sites in each county which provide fee and unbiased information about Medicare. For more information, go to [http://www.ncdoi.com/shiip](http://www.ncdoi.com/shiip).

**Project C.A.R.E.** (Caregiver Alternatives to Running on Empty) is a state funded dementia-specific support program for family caregivers. The program uses family consultants to provide support to caregivers caring for someone with Alzheimer’s disease or related dementia at home. Family consultants assist caregivers with education, resources and access to respite vouchers to relieve some of the stress caregiving can create. The level of assistance provided depends on what the caregiver needs and prefers. For more information about accessing Project C.A.R.E. in your region, go to [https://www.ncdhhs.gov/assistance/adult-services/project-care](https://www.ncdhhs.gov/assistance/adult-services/project-care), or contact Dawn Oakey Gartman, State Project C.A.R.E. Director, at 919-855-3462.

**Project Lifesaver** is a free rapid response program that locates children or adults with brain disorders who wander away from their caregivers. Individuals enrolled in the program wear a bracelet that contains a battery-operated transmitter that emits a tracking signal. Contact your local sheriff’s office for more information.

**Silver Alert** is an alert system to notify the public about missing individuals with dementia or other cognitive impairments so that they can be found quickly and reunited with their caregivers. Contact local law enforcement to report the missing person. There is no minimum length of time before a report can be made.

**VA Caregiver Support Program** provides a variety of support services for family caregivers of Veterans. To learn more about what the program offers, call 1-855-260-3274 or go to [https://www.caregiver.va.gov/support/support_services.asp](https://www.caregiver.va.gov/support/support_services.asp).

*July 1, 2017*